Corps of Engineers, Department of the Army

33 CFR Part 209

Administrative Procedures; Removal of Obsolete Requirements; Correction

AGENCY: Army Corps of Engineers, DoD. **ACTION:** Final rule: Correction.

SUMMARY: The regulations which prescribe the procedures used by the Corps of Engineers in the review and promulgation of regulations governing navigable waters were amended to remove obsolete and unnecessary requirements on December 22, 1986, 51 FR 45765. An error made in the designation of paragraphs in 33 CFR 209.200 in hereby corrected.

EFFECTIVE DATE: June 29, 1987.

ADDRESS: USACE, DAEN-CECW-OR, Washington, DC 20314-1000.

FOR FURTHER INFORMATION CONTACT: Mr. Ralph T. Eppard or Mr. Sam Collinson at (202) 272–1783.

SUPPLEMENTARY INFORMATION: The amendatory language to \$ 209.200 as it appeared in 51 FR 45765, December 22, 1986, is corrected as follows:

5. Section 209.200 Regulations governing navigable waters is amended by revising paragraph (a), removing paragraphs (d) and (f) and redesignating paragraph (e) as (d), paragraph (g) as (e) and redesignating paragraph (h) as (f), as follows:

Dated: June 22, 1987.

John O. Roach, II.

Army Liaison Officer With the Federal Register.

[FR Doc. 87-14520 Filed 6-26-87; 8:45 am] BILLING CODE 3710-92-M

ENVIRONMENTAL PROTECTION AGENCY

40 CFR Parts 795 and 799

[OPTS-42065B; FRL-3223-9]

2-Ethylhexanoic Acid; Technical Modification

AGENCY: Environmental Protection Agency (EPA).

ACTION: Final rule.

SUMMARY: EPA has approved minor modifications of the test standards for the 2-ethylhexanoic acid (EHA) test rule at 40 CFR 795.223(c)(2)(ii)(C) and 799.1650 (c)(2)(ii)(C) and (c)(3) in response to a request from the test sponsors.

EFFECTIVE DATE: June 29, 1987.

FOR FURTHER INFORMATION CONTACT: Edward A. Klein, Director, TSCA Assistance Office (TS-799), Office of Toxic Substances, Rm. E-543, 401 M St. SW., Washington, DC. 20460, (202) 554-

SUPPLEMENTARY INFORMATION: EPA has approved minor modifications of the test standards for the 2-ethylhexanoic acid test rule at 40 CFR 795.223 and 799.1650.

Test Rule Modifications

EPA issued a final test rule for EHA. published in the Federal Register of November 6, 1986 (51 FR 40318). On March 19, 1987, the Chemical Manufacturers Association (CMA) submitted a request to remove the requirement that the subchronic toxicity test be administered by the same route as the pharmacokinetics test. The Agency reviewed the request and has decided to grant the requested modification. Because the modification merely clarifies an inconsistency, the minor changes to 40 CFR 795.223(c) (2) (ii)(C) and 799.1650 (c)(2)(ii)(C) and (c)(3) requested by CMA clearly do not pose any substantive issues. Moreover, the Agency has determined that obtaining comment on these modifications would be impracticable, unnecessary, and contrary to the public interest in that doing so would delay both the start of testing and the submission of the study data. Therefore, in accordance with the procedures in 40 CFR 790.55(b)(1)(ii) and section 553(b)(3)(B) of the Administrative Procedure Act, the Agency notified the test sponsors by letter of its approval of these modifications on April 7, 1987 without seeking public comment (Ref. 2). In accordance with 40 CFR 790.55(b)(2), EPA is issuing this notice which describes the associated modifications to the test standards and reporting requirements for publication in the Federal Register. For a more detailed description of the rationale for these modifications, refer to CMA's letter (Ref. 1) and EPA's letter in response (Ref. 2). The rule is modified as follows:

The EHA rule requires an oral subchronic toxicity study to develop data comparable with other data on EHA and data on related chemicals like 2-ethylhexanol and di(2-ethylhexyl) phthalate. To accomplish this, the Agency believes it is unnecessary to restrict the EHA oral subchronic toxicity test to gavage and encapsulation as required for the pharmacokinetics test (Ref. 2). Furthermore, the test sponsors have conducted a preliminary stability study (Ref. 3) that indicates a traditional feeding study may be technically feasible. Therefore, the Agency has decided to remove the requirement in

the final rule that both the pharmacokinetics and subchronic toxicity tests utilize either gavage or encapsulation.

The 3-month extension in the final reporting requirement for the oral subchronic toxicity test allows the test sponsors adequate time to complete validation studies for alternative oral routes of administration and to complete testing using a fully validated oral route of administration.

Public Record

EPA has established a public record for this rulemaking [docket number OPTS-42065B]. The record includes the information considered by the Agency in evaluating the requested modifications to this rule.

(1) The Chemical Manufacturers Association (CMA). Letter from Geraldine Cox, Ph.D., to the Director. Office of Compliance Monitoring, Office of Pesticides and Toxic Substances, USEPA. (March 19, 1987).

(2) USEPA. Letter from Charles E. Elkins, Director, Office of Toxic Substances, to Geraldine Cox. CMA. (April 7, 1987).

Geraldine Cox, CMA. (April 7, 1987).
(3) Eastman Kodak Co. Dose verification,
Analysis, and Recovery of 2-Ethylhexanoic
Acid in Rat Chow, prepared for the EHA
Panel of CMA, Washington, DC 20037
(December 19, 1986).

The record is available for inspection from 8 a.m. to 4 p.m., Monday through Friday, except legal holidays, in Rm. G-004, NE Mall, 401 M St., SW., Washington, DC 20460.

Dated: June 17, 1987.

John A. Moore,

Assistant Administrator for Pesticides and Toxic Substances.

Therefore, 40 CFR Chapter I is amended as follows:

PART 795—[AMENDED]

- 1. In Part 795:
- a. The authority citation continues to read as follows:

Authority: 15 U.S.C. 2603, 2625.

b. In § 795.223 by revising paragraph (c)(2)(ii)(C), to read as follows:

§ 795.223 Pharmacokinetics test.

- (c) * * *
- (2) * * *
- . iii * * *
- (C) Oral dosing shall be performed by gavage or by administering encapsulated test substance.

Part 799—[AMENDED]

2. In Part 799:

a. The authority citation for Part 799 continues to read as follows:

Authority: 15 U.S.C. 2603, 2611, 2625. b. In § 799.1650 by revising paragraphs (c)(2)(ii)(C) and (c)(3) to read as follows:

§ 799.1650 2-Ethylhexanoic acid.

- (c) * * *
- (2) * * * (ii) * * *
- (C) The final report of results shall be submitted to the Agency no later than 18 months from the effective date of the final test rule.
- (3) Administration of test substance. Dosing for the testing required under paragraph (c) (1) and (2) of this section shall be by the oral route for both tests, and as specified in § 795.233(c)(2)(ii)(C) and § 795.260(d)(7) of this chapter

[FR Doc. 87–14673 Filed 6–28–87; 8:45 am]

GENERAL SERVICES ADMINISTRATION

41 CFR Part 101-20

[FPMR Amdt D-84]

Management of Buildings and Grounds

AGENCY: General Services Administration.

ACTION: Final rule; Correction.

SUMMARY: General Services
Administration is correcting minor,
nonsubstantive errors to the final rule,
FPMR Amendment D-84, which governs
the operation of, and the activities in
Federal buildings. This regulation was
developed by a subcommittee of the
Interagency Advisory Committee on
Regulatory Review and is designed to
clarify the content, eliminate
duplication, and remove obstacles to
effective building management.

EFFECTIVE DATE: April 8, 1987.

FOR FURTHER INFORMATION CONTACT: James Steele (202-566-1563).

SUPPLEMENTARY INFORMATION: In FR document 87–7694 appearing at 52 FR 11263, Apr. 8, 1987, GSA revised Part 101–20. This document corrects several incorrect internal references.

Dated: June 23, 1987.

Rodney P. Lantier,

Chief, Directives and Reports Managemem Branch.

PART 101-20—[AMENDED]

Therefore, 41 CFR Part 101-20 is corrected as follows:

§ 101-20.103-4 [Corrected]

On page 11267, first column, in paragraph (a) the reference reading "§ 101-20.003-7" is corrected to read "§ 101-20.003(g)." In the second column, in paragraph (a) the reference reading "§ 101-20.003-23" is corrected to read "§ 101-20.003(w)." In paragraph (c) the reference reading "§101-20.003(w)." In paragraph (c) the reference reading "§ 101-20.003(x)." In paragraph (d) the reference reading "§ 101-20.003-22" is corrected to read "§ 101-20.003(v)."

§ 101-20.107 [Corrected]

On page 11270, third column, the section title reading "Energy management" is corrected to read "Energy conservation." On page 11271, second column, in paragraph (h) the reference reading "\square 101-20.116" is corrected to read "\square 101-20.107."

§ 101-20.302 [Corrected]

On page 11273, first column, the reference reading "§ 101-20.003-7" is corrected to read "§ 101-20.003(g)."

§ 101-20.309 [Corrected]

On page 11273, third column, the reference reading "\\$ 101-20.003-28" is corrected to read "\\$ 101-20.003(z)."

§ 101-20.403 [Corrected]

On page 11274, third column, in paragraph (a)(2) the reference reading "§ 101-20.003-4" is corrected to read "§ 101-20.003(d)."

§ 101-20.404 [Corrected]

On page 11274, third column, in paragraph (a) the reference reading "§ 101-20.003-13" is corrected to read "§ 101-20.003(m)."

[FR Doc. 87-14634 Filed 6-26-87; 8:45 am]
BILLING CODE 6820-22-M

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Public Health Service

42 CFR Part 57

Programs for the Training of Physician Assistants and Grants for Physician Assistant Training Programs

AGENCY: Public Health Service, HHS. **ACTION:** Final regulations.

SUMMARY: These final regulations: (1)
Amend the current regulatory definition
of programs for the training of physician
assistants in accordance with section
701(8) of the Public Health Service Act
(the Act) (42 CFR Part 57, Subpart I), as
amended by the Health Professions
Training Assistance Act of 1985 (Pub. L.

99–129); and (2) amend the existing regulations governing the Grants for Physician Assistant Training Programs, authorized by section 783 of the Act (42 CFR Part 57, Subpart H) to: (a) Conform the regulations with amendments made to the Act by Pub. L. 99–129, the Omnibus Budget Reconciliation Act of 1981 (Pub. L. 97–35), and the Compact of Free Association Act of 1985 (Pub. L. 99–239), which are of a technical and ministerial nature; and (b) add provisions to encourage efforts to attract, maintain and graduate minority and disadvantaged students.

EFFECTIVE DATE: These regulations are effective June 29, 1987.

FOR FURTHER INFORMATION CONTACT:

Donald L. Weaver, M.D. Director, Divsion of Medicine, Bureau of Health Professions, Health Resources and Services Administration, Parklawn Building, Room 4C25, 5600 Fishers Lane, Rockville, Maryland 20857; telephone: 301 443–6190.

SUPPLEMENTARY INFORMATION: On October 10, 1986, the Assistant Secretary for Health, Department of Health and Human Services, with the approval of the Secretary, published in the Federal Register (51 FR 36412), a Notice of Proposed Rulemaking (NPRM) to amend Subpart I of 42 CFR Part 57 to implement changes in section 701(8) of the Act. Pub. L. 99-129, amended section 701(8) by: (a) Substituting the term "primary health" for "health" to describe the type of care that graduates of physician assistant training programs must be capable of providing under the supervision of a physician; and (b) adding a requirement that physician assistant programs train students in primary care, disease prevention, health promotion, geriatric medicine, and home health care. The statute further directs the Secretary to consult with appropriate organizations and then promulgate regulations to implement the amendments. The American Medical Association, American Academy of Physician Assistants and the Association of Physician Assistant Programs provided comments to the Public Health Service in the development of the proposed new

In the NPRM of October 10, 1986 (51 FR 36414), the Department also proposed to amend the existing regulations governing grants under section 783 of the Act, codified at 42 CFR Part 57. Subpart H. This amendment is necessary to provide an added emphasis on the national need to train more minority and disadvantaged students.

definitions.